## Amendments to the Claims:

This listing of claims will replace all prior versions, and listings, of claims in the application:

# **Listing of Claims:**

 (Currently Amended) A computer-implemented method for displaying information on one or more user interfaces regarding the likelihood a person has a gene variant indicative of an atypical event, comprising the steps of:

> displaying a first user interface to a clinician, the user interface configured to display and receive clinical agent information including at least one identifier of a clinical agent;

> receiving from the user interface the clinician's inputs including at least one identifier of a clinical agent and a dosage of the clinical agent, wherein receiving includes receiving a selection of an entry in a listing of clinical agents on the first user interface and a selection of the dosage from a range of dosages recommended for the clinical agent associated with the selected entry;

accessing a data structure to determine if a gene variant is known to be associated with one or more atypical events for the identifier of the clinical agent received from the clinician, wherein the data structure includes an agent-gene association table information;

inquiring if the person to whom the clinical agent is to be administered has a stored genetic test result value for the gene variant, wherein inquiring includes accessing an electronic medical record (EMR) of the person;

accessing hereditary information for the person if the person does not have a genetic test result value for the genetic variant, the hereditary information being

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information that may be utilized to determine if the person has a predisposition for certain conditions, wherein the hereditary information is obtained from the EMR of the person;

determining from the hereditary information whether a parent of the person had the gene-variant;

utilizing the hereditary information for the person to determine the likelihood the person has the gene variant;

generating an output including information regarding the likelihood that the person has the gene variant indicative of an atypical event based on the hereditary information; and

displaying a second user interface to the clinician, the user interface configured to display the output regarding the likelihood the person has the gene variant indicative of an atypical event for the identifier of the clinical agent received from the clinician based on the hereditary information.

## (Canceled).

 (Previously Presented) The method of claim 1, wherein the hereditary information includes ethnicity, gender or geographic origin.

#### (Canceled)

5. (Previously Presented) The method of claim 1, wherein the accessing of the hereditary information comprises accessing the hereditary information from an electronic medical record of the person stored within a comprehensive healthcare system.

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(Previously Presented) The method of claim 1, further comprising

the step of initiating one or more clinical actions if a test result value is not available for the

person and the information regarding the risks indicates a significant risk that the person has the

gene variant associated with an atypical event.

7. (Previously Presented) The method of claim 6, wherein the one or

more clinical actions comprise at least one of ordering a genetic test, and displaying a third user

interface configured to display a notification to the clinician that the agent should be suspended

in view of the risk of an atypical event, canceling the order for the clinical agent absent input

from the clinician to the contrary, and recommending an alternative clinical agent.

8. (Currently Amended) A computer system embodied on one or more

computer storage media having computer-executable instructions embodied thereon for

displaying information on one or more user interfaces regarding the likelihood that the person

has the gene variant indicative of an atypical event based on the hereditary information,

comprising:

a first displaying component that displays a first user interface to a

clinician, the user interface configured to display and receive clinical agent

information including at least one identifier of a clinical agent;

a receiving component that receives from the user interface the clinician's

inputs including at least one identifier of a clinical agent and a dosage of the

clinical agent, wherein receiving includes receiving a selection of an entry in a

listing of clinical agents on the first user interface and a selection of the dosage

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from a range of dosages recommended for the clinical agent associated with the

selected entry;

a first accessing component for accessing a data structure to determine if a

gene variant is known to be associated with one or more atypical events for the

identifier of the clinical agent received from the clinician, wherein the data

structure includes an agent-gene association table information:

an inquiring component that inquires if the person to whom the clinical

agent is to be administered has a stored genetic test result value for the gene

variant, wherein inquiring includes accessing an electronic medical record (EMR)

of the person;

a second accessing component for accessing hereditary information for the

person if the person does not have a genetic test result value for the gene variant,

the hereditary information being information that may be utilized to determine if

the person has a predisposition for certain conditions, wherein the hereditary

information is obtained from the EMR of the person;

a determining component for determining from the hereditary information

whether a parent of the person had the gene variant;

a utilizing component for utilizing the hereditary information for the

person to determine the likelihood the person has the gene variant;

a generating component that generates an output including information

regarding the likelihood that the person has the gene variant indicative of an

atypical event based on the hereditary information; and

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a second displaying component for displaying a second user interface to the clinician, the user interface configured to display the output regarding the likelihood the person has the gene variant indicative of an atypical event for the identifier of the clinical agent received from the clinician based on the hereditary information.

## 9. (Canceled)

 (Previously Presented) The computer system of claim 8, wherein the hereditary information includes ethnicity, gender or geographic origin.

#### 11. (Canceled)

12. (Previously Presented) The computer system of claim 8, wherein the second accessing component accesses the hereditary information from an electronic medical record of the person stored within a comprehensive healthcare system.

13. (Previously Presented) The computer system of claim 8, further comprising an initiating component that initiates one or more clinical actions if a test result value is not available for the person and the information regarding the risks indicates a significant risk that the person has the gene variant associated with an atypical event.

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14. (Previously Presented) The computer system of claim 13, wherein

the one or more clinical actions comprise at least one of ordering a genetic test and displaying a

third user interface configured to display a notification to the clinician that the agent should be

suspended in view of the risk of an atypical event, canceling the order for the clinical agent

absent input from the clinician to the contrary, and recommending an alternative clinical agent,

15. (Currently Amended) A computer storage medium containing instructions

for a method for controlling a computer system for displaying information on one or more user

interfaces regarding the likelihood that the person has the gene variant indicative of an atypical

event based on the hereditary information, the method comprising the steps of:

displaying a first user interface to a clinician, the user interface configured

to display and receive clinical agent information including at least one identifier

of a clinical agent;

receiving from the user interface the clinician's inputs including at least

one identifier of a clinical agent and a dosage of the clinical agent, wherein

receiving includes receiving a selection of an entry in a listing of clinical agents

on the first user interface and a selection of the dosage from a range of dosages

recommended for the clinical agent associated with the selected entry;

accessing a data structure to determine if a gene variant is known to be

associated with one or more atypical events for the identifier of the clinical agent

received from the clinician, wherein the data structure includes an agent-gene

association table information;

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inquiring if the person to whom the clinical agent is to be administered has a stored genetic test result value for the gene variant, wherein inquiring includes accessing an electronic medical record (EMR) of the person;

accessing hereditary information for the person if the person does not have a genetic test result value for the genetic variant, the hereditary information being information that may be utilized to determine if the person has a predisposition for certain conditions;

determining from the hereditary information whether a parent of the person had the gene variant:

utilizing the hereditary information for the person to determine the likelihood the person has the gene variant;

generating an output including information regarding the likelihood that the person has the gene variant indicative of an atypical event based on the hereditary information; and

displaying a second user interface to the clinician, the user interface configured to display the output regarding the likelihood the person has the gene variant indicative of an atypical event <u>for the identifier of the clinical agent received from the clinician based on the hereditary information</u>.

## (Canceled)

17. (Previously Presented) The computer storage medium of claim 15, wherein the hereditary information includes ethnicity, whether the person's parents had the gene variant, gender and geographic origin.

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18. (Canceled)

(Previously Presented) The computer storage medium of claim 15,

wherein the accessing of the hereditary information comprises accessing the hereditary information from an electronic medical record of the person stored within a comprehensive

healthcare system.

20. (Previously Presented) The computer storage medium of claim 15,

further comprising the step of initiating one or more clinical actions if a test result value is not

available for the person and the information regarding the risks indicates a significant risk that

the person has the gene variant associated with an atypical event.

21. (Previously Presented) The computer storage medium of claim 20,

wherein the one or more clinical actions comprise at least one of ordering a genetic test and

displaying a third user interface configured to display a notification to the clinician that the agent

should be suspended in view of the risk of an atypical event, canceling the order for the clinical

agent absent input from the clinician to the contrary, and recommending an alternative clinical

agent.

22. (Previously Presented) The method of claim 1, wherein the hereditary

information comprises one of the genetic characteristics of the person's family members or other

relevant family history, demographic information for the person or combinations thereof.

wherein the demographic information is indicative of genetic predisposition to certain

conditions.

23. (Previously Presented) The method of claim 1, further comprising:

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> storing the output including information regarding the likelihood that the person has a gene variant indicative of an atypical event based on the hereditary information in the patient's electronic medical record.

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